(11) Publication number:

0 101 878

A₂

12)

EUROPEAN PATENT APPLICATION

(21) Application number: 83107026.3

(22) Date of filing: 18.07.83

(5) Int. Ci.³: **C** 10 **G** 11/18 **C** 10 **G** 57/00

(30) Priority: 29.07.82 US 403241

43 Date of publication of application: 07.03.84 Bulletin 84/10

(B4) Designated Contracting States: AT BE DE FR GB IT NL SE

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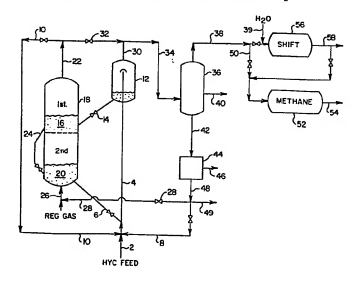
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(54) Combination process for upgrading reduced crude.

(5) In a reduced crude conversion the catalyst is regener- combined with hydrogen to produce methane, methanol or to CO and H₂ with minimum CO₂ formation. The CO can be produce addition H₂.

ated with mixture of oxygen-enriched gas and H₂O either as Fischer-Tropsch liquid hydrocarbons or can be passed steam or preferably as water to provide additional expansion through the reduced crude conversion operation as lift gas, and facilitate fluidization, while converting coke on catalyst or as cooling medium or subjected to water-gas shift to



A COMBINATION PROCESS FOR UPGRADING REDUCED CRUDE

Background of the Invention

- The combination operation of this innovative processing sequence embodies scattered excerpts of the prior art in a synergistic relationship particularly contributing to efficient reduced crude processing.
- The reaction of steam with coke on solid substrates was the subject of a paper by T.Y. Yan and M.P. Rosynek given before the American Chemical Society in September 1979, Washington Meeting. In this paper reference is made to the catalytic processing of heavy oil and residua over cracking catalysts and various schemes for removing deposited coke using steam and oxygen by referring to U.S. Patents 3691063, 3726791 and 3983030.
- 20 U.S. Patent 3433732 discloses steam reforming a heated hydrocracking catalyst to produce hydrogen and regenerate the catalyst at the same time.
- U.S. Patent 2888395 discloses a catalytic coking process whereby coked catalyst is steam reformed to produce hydrogen.
- U.S. Patent 2702267 discloses the mixing of spent and regenerated catalyst particles in a soaking zone and using hot regeneration product gases as the fluidizing and stripping gas in the soaking zone. The catalyst is regenerated at a temperature above 760°C (1400°F) with a mixture of steam and high purity oxygen comprising no more than 10 vol. % of nitrogen to produce a flue gas product comprising hydrogen, carbon oxides and

excess steam. When the mixture of steam and high purity oxygen is used as the regenerating medium, the principal reactions are:

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$$2^{C+O_2-2CO}$$
 (Exoth) $C+H_2O-CO+H_2$ (Endo) $C+O_2-CO_2$ (Exoth) $C+CO_2-2CO$ (Endo)

To insure a high yield of hydrogen, more steam than oxygen is used.

British Patent 2001545 describes a two stage regeneration operation controlled so that only partial regeneration of catalyst particles occurs in the first zone and the catalyst particles are not heated excessively and a CO flue gas is produced.

The use of CO₂ or steam with an oxygen containing steam is disclosed whereby a flue gas of high CO or CO-hydrogen concentration is obtained. Such a steam is valuable for use in water-gas shift to produce hydrogen. The regeneration zones are positioned one above the other so that partially regenerated catalyst may flow from the upper zone to the lower zone by gravity. Regeneration gas used in upper zone may be either (2+steam, O2+CO2, O2+steam+CO2 depending on composition of flue gas particularly desired. Preferably the gas charged to the second stage of regeneration if air.

U.S. Patent 4244811 discloses the catalytic cracking of a hydrocarbon feed in the presence of water and subjecting deactivated catalyst with coke deposits to gasification conditions consisting of partial oxidative combustion to produce gas rich in CO or with the addition

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of steam to produce a gas rich in hydrogen or both CO + H_2 .

U.S. Patent 3691063 relates to a residual fuel oil hydrocracking process wherein a residuum feed is subjected to metals removal in a guard case containing an acid catalyst along with asphaltenes. The guard catalyst is regenerated with steam and oxygen to maximize hydrogen production by partial oxidation of asphaltenes. The hydrogen produced is used in the hydrocracking step. Hydrogen plus carbon monoxide from the guard chamber regeneration is fed to a two stage water gas shift operation where steam is reacted with the carbon monoxide to form additional hydrogen and carbon dioxide.

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U.S. Patent 3726791 related to a process where high Conradson carbon feeds are coked to lay down carbon deposits on a gasification catalyst and the coked catalyst is steam gasified to produce hydrogen.

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U.S. Patent 3983030 relates to a process for demetalation and desulfurization of residua and deposited coke gasification with steam and free oxygen to produce producer gas and regenerate a porous refractory oxide used to demetallize and desulfurize the residual.

Summary of the Invention

In processing a reduced crude feed or a residual oil boiling above about 343°C (650°F) comprising metal contaminants and Conradson carbon producing components, there are several significant factors one must consider and deal with in the operation. These

factors include large deposition of hydrocarbonaceous material, more often referred to as coke or coke make on the catalyst, metals contaminant level, and the presence of sulfur and to a lesser extent nitrogen. The amount of coke made is a function of feed composition and severity The amount of of the hydrocarbon conversion operation. coke made also influences the regeneration operation at least with respect to the temperatures encountered to obtain desired coke removal and optimize regenerated catalyst activity, since maximum catalyst activity and selectivity should result from eliminating coke or carbon residue on the catalyst. In addition to the above it is necessary to control the deactivating effects of metal to minimize undesired hydrogen contaminants so as of desired product at expense the production increased coke deposition particularly production of promoted by reduced metals on the catalyst. important factor requiring consideration is particularly concerned with controlling the emissions of sulfur oxides and nitrogen oxides or their recovery from the processing unit and particularly the regeneration operation of the process.

In the combination process of this invention it is proposed in one aspect that the spent catalyst of hydrocarbon conversion comprising carbonaceous deposits and containing hydrogen by regenerated with a mixture of oxygen rich gas greater than provided by air and water such as steam so as to facilitate the fluid catalyst regeneration while converting deposited carbonaceous material (coke) to particularly CO with the oxygen-steam mixture under conditions to restrict production of CO₂ to a low minimum. Thus it is proposed to realize a flue gas

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product of the regeneration operation comprising CO, sulfur oxides, nitrogen oxides, steam and hydrogen and some CO₂ absent combustion supporting amounts of oxygen considerably enhanced regeneration temperature In combination with the above the flue gas thus produced is cooled and condensed to remove water, sulfur oxides and nitrogen oxides leaving a CO rich flue gas comprising some CO₂ and hydrogen. In one particular embodiment it is contemplated combining the CO rich flue gas of regeneration above recovered with a hydrogen rich gas product stream of an adjacent hydrocarbon conversion operation to provide a syngas product convertable to methane and/or methanol for use as herein discussed. yet another embodiment, the CO rich flue gas above recovered is charged with the high boiling hydrocarbon feed as a fluidizing and atomizing gas to the reaction zone and to help with obtaining intimate contact with particles of catalyst in combination with reducing coke and hydrogen production as herein discussed.

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The use of an oxygen enriched regeneration gas or one of high oxygen purity at least twice that of air comprising steam is instrumental in recovering a CO rich flue gas free of nitrogen oxides from the catalyst regeneration and combination operation of the invention. The recovered CO rich regeneration flue gases of lower CO₂ content may also be used for stripping catalyst separated from vaporous hydrocarbon products and before affecting regeneration of the stripped catalyst by the method of the invention. The hydrocarbon conversion-regeneration technique of this invention permits controlling regeneration temperatures within a relatively narrow range during reduction of deposited

coke to a desired low level below 0.25 weight percent and preferably below 0.1 weight percent, reduced metals activity in the conversion zone, improves product selectivity of the hydrocarbon conversion operation, and considerably reduces the use of expensive air compressors for achieving adequate fluidization of catalyst particles during regeneration thereof.

The processing combination of this invention the combination synergistics relationship of 10 efficient-economical more provide а operation to operation is particularly geared to processing topped crudes, reduced crudes, vacuum bottoms, heavy asphaltic crudes, coal crude oils, shale oils and tar sands oil products. The fluid catalystic cracking operation of the 15 invention particularly combination process of this comprises an elongated riser reactor discharging into a catalyst-hydrocarbon vapor separating vessel and catalyst The catalyst is separated from collecting vessel. hydrocarbon product vapors upon discharge from the riser 20 as by a ballistic separation technique or other suitable techniques known in the industry. The separated catalyst is collected and passed to a catalyst stripping zone and thence is passed to catalyst regeneration comprising in particular a two stage catalyst regenerated operation 25 herein discussed. A regeneration vessel or apparatus arrangement such as provided in copending application filed 4/19/82 or that of 6026 10/21/82 may be employed by the method of this invention. In any one of these apparatus arrangements 30 the spent catalyst comprising hydrocarbonaceous deposits and referred to herein as coke is charged to an upper fluid bed of catalyst particles being regenerated as

herein provided. In this first stage of regeneration the spent catalyst is partially regenerated at temperatures below 760°C (1400°F) by removing from 40 to 80% of the coke deposits on the catalyst in the presence of partially spent oxygen rich regeneration gases comprising 5 CO₂ recovered from a lower regeneration section as herein discussed. The partially regenerated catalyst thus obtained is then passed by gravity from the upper bed of catalyst in the first regeneration zone to a lower bed of catalyst in a second regeneration zone where it contacts 10 fresh high purity oxygen to steam or $^{\rm O}_2$ plus $^{\rm CO}_2$ regeneration gases or other high purity oxygen rich gas to produce a regenerated catalyst comprising residual carbon less than about 0.1 weight percent and preferably 15 no more than about 0.05 weight percent residual carbon. Catalyst particles thus regenerated and recovered at a temperature below 760°C (1400°F) are passed to the riser cracking zone wherein the catalyst forms a suspension with a mixture of hydrocarbon feed, water and with or without a CO rich fluidizing gas obtained as herein 20 provided to initiate a new cycle of the combination process.

The fluidizing gas particularly employed in the riser reactor of the combination operation of this 25 invention is a particular regeneration flue gas product produced as herein provided and exiting from the two stage regeneration operation described. The make up of this flue product gas is controlled oxygen-steam- CO_2 content of the regeneration gas employed 30 in the combination regeneration operation. That is the make up of the flue gas is determined by the steam-oxygen content or ratio of the regeneration gas initially

charged to the bottom regeneration section and the temperature control exercised upon the dilute and dense catalyst phase thereof. For example, the use of air and a regeneration temperature equal to or less than 732°C (1350°F) will yield a CO-nitrogen containing flue gas of 5 low ${\rm CO}_2$ content. As temperatures are raised through the use of excess oxygen or CO burning in the dense or dilute catalyst phases, the flue gas will comprise CO_{2} unconsumed oxygen and less nitrogen as regeneration air is enriched with oxygen. Thus the nitrogen content of 10 the flue gas may be considerably reduced by using oxygen enriched air or a high purity oxygen-CO2 rich stream as the regeneration gas to produce CO and/or CO2 rich flue gases of low or no nitrogen oxides. Since sulfur oxides are a product of the hydrocarbon feed, little can be done 15 except as provided herein to further reduce sulfur oxides in flue gas.

The use of a CO and/or CO_2 containing flue gas product of the regeneration operation as a fluidizing gas 20 in the hydrocarbon riser reaction zone is of considerable advantage and may be used to control the hydrogen produced by catalytic conversion of the heavy hydrocarbon feed by particularly keeping the metals in the feed and deposited on the catalyst in an oxidized state which also 25 operates to reduce the formation of carbon by reduced This particular fluidizing gas composition comprising CO and ${\rm CO}_2$ will also undergo to some extent the well known water gas shift reaction with water introduced with the feed at the hydrocarbon conversion 30 Hydrogen produced by the water gas shift conditions. reaction will operate to suppress hydrogen produced by metals cracking without substantially interferring with

suppression of coke formation. The CO-steam containing flue gas recovered as high regeneration temperatures up to 760°C (1400°F) may also be used as a heat sink source to supplement the endothermic heat of reaction required in the hydrocarbon riser conversion zone. This CO-steam 5 rich flue gas may be introduced to the bottom of the riser or at one or more points along the length of the In addition the CO-steam rich flue gas may be introduced to the bottom of the riser or at one or more points along the length of the riser. 10 In addition the CO-steam containing flue gas of relatively low CO2 content may be introduced to the collected catalyst following separation of hydrocarbon vapors to effect an initial high temperature stripping of the separated catalyst.

A CO_2 rich flue gas may be employed to strip regenerated catalyst initially contacted with high purity oxygen prior to passage thereof to the hydrocarbon conversion zone to remove oxygen from the catalyst.

In one particular embodiment, is contemplated mixing a CO rich gas product of catalyst regeneration with or without formed hydrogen therein with a hydrogen rich gaseous product recovered from the vaporous products of hydrocarbon conversion to form a syngas mixture thereof. In this particular embodiment, light gaseous product hydrocarbons recovered vaporous products of hydrocarbon conversion are treated to removed formed ammonia and acid components such as sulfur oxides, hydrogen sulfide to provide a light hydrocarbon stream comprising hydrogen. The hydrogen rich gases thus recovered are mixed with CO rich gases of

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the regeneration operation to form a syngas mixture charged to a methanation reactor zone, or a methanol conversion unit, to form methane and/or methanol. The exothermic reaction heat thus generated is recovered by heat exchange means not shown and/or are used with produced methane to furnish heat to the regeneration operation and/or used as fuel to furnish power to drive a gas turbine to generate compressor power.

The combination process of this invention thus 10 contemplates the several variations herein identified and used either alone or in combination with one another to achieve the following particular operating concepts. That is, an oxygen enriched gas in combination with steam is used to regenerate catalyst under conditions to 15 produce a CO rich flue gas containing hydrogen in combination with little CO2 and very little nitrogen oxide and sulfur oxide containing materials. gas thus produced is used either with or without sulfur oxides removal as fluidizing gas in a reduced crude riser 20 conversion zone or a portion thereof is mixed with hydrogen rich product gases of the hydrocarbon conversion operation with or without gas shift, and manipulated to achieve additional hydrogen production than contributed by the hydrocarbon feed within the riser cracking zone. syngas mixture thus obtained is subjected to a 25 further water gas shift operation in one embodiment to produce addition syngas baefore converting the syngas product of the operation to methanation or methanol temperature of exothermic The production. 30 methanation reaction may be used to preheat gases charged to the regeneration operation and/or provide fuel for a

fired turbine prime mover arrangement providing compressed regeneration oxygen containing gases.

In yet another embodiment, it is particularly 5 contemplated stripping spent catalyst comprising carbonaceous deposits before oxygen combustion in admixture with hot regenerated catalyst particles to provide a mix temperature thereof within the range of 566°C (1050°F) to about 649°C (1200°F) and using as a fluidizing and stripping medium, flue gas products of 10 regeneration of selected CO and/or CO2 content obtained as provided herein. Thus the stripping medium may be selected from a number of gasoue product materials of the process comprising steam, CO, CO2, and mixtures thereof 15 particularly contributing to controlled temperature reaction of spent hydrocarbon conversion catalyst. this operating environment the catalyst may be initially subjected to steam stripping at the temperature recovered the hydrocarbon conversion operation and then further stripped at the higher temperature in admixture 20 hot regenerated catalyst particles discussed. The first stage οf stripping may accomplished in a dense fluid bed of catalyst external to or about the riser reactor and the second stage stripping may be accomplished in an upflowing riser contact zone or 25 dense fluid bed contact zone under conditions particularly encouraging the removal of insufficiently converted hydrocarbon material from the catalyst with CO, CO2, steam and combinations thereof suitable for the 30 Catalyst thus stripped is then passed to the purpose. first stage of catalyst regeneration of the two stage regeneration operation herein discussed. In this two stage regeneration, oxygen enriched air or high purity

oxygen containing gas admixed with considerable steam is charged to the second stage of regeneration in an amount sufficient to achieve substantially total removal of deposited carbonaceous material from the catalyst as charged to the first stage of regeneration and product gases of the second stage regeneration operation are charged to the first regeneration stage to accomplish up to 80% removal of deposited carbonaceous material therein and produce a flue gas stream absent combustion supporting amounts of oxygen particularly comprising a CO rich flue gas stream containing steam.

Discussion of Specific Embodiments

15 The drawing is a diagrammatic sketch in elevation of the combination process of this invention as particularly related to the catalytic conversion of reduced crudes, regeneration of catalyst used therein to produce of CO rich flue gas comprising steam, the recovery of hydrogen rich gases from the products of a selective reduced crude hydrocarbon conversion operation and the generation of methane and/or methanol heat for use in the process from syngas products of catalyst regeneration and reduced crude conversion.

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Referring now to the drawing by way of example there is shown a diagrammatic arrangement of interconnected vessels or process zones for practicing the concepts of this invention particularly directed to processing reduced crudes. In the arrangement of the drawing, a reduced crude feed or other high boiling material herein identified is charged by conduit 2 to a riser reaction zone 4 wherein it contacts hot regenerated

catalyst charged to riser 4 by standpipe 6. In addition thereto a condensed water product of the process which has been treated to partially remove sulfur and nitrogen compounds therein is charged for admixture with the feed 5 the other hand water with conduit 8. On dissolved sulfur compound may be charged with advantage the riser to suppress undesired deposited metal contaminant activity. To facilitate contact of the heavy oil feed with the catalyst as herein provided 10 discussed above, а special flue gas product of regeneration obtained as herein provided may also be charged for admixture with the hydrocarbon feed by It is to be understood that the materials conduit 10. charged by conduits 10, 8 and 2 do not necessarily need to meet at a common point but may be charged in a manner 15 most expedient to achieve mixing and atomization of the charged high boiling hydrocarbon feed for dispersed phase contact with hot regenerated catalyst particles. conversion of the hydrocarbon feed in riser 4 employing 20 equilibrium conversion catalyst comprising metal contaminant deposits above fresh feed catalyst up to about 30,000 ppmNi equivalent is monitored and controlled to achieve selective conversion hydrocarbon feed in the presence of hydrogen generated in the operation as discussed above in a reaction time frame 25 restricting undesired reactions promoted by reduced metal contaminants. Thus in the particular hydrocarbon conversion operation of this invention the reactants charged to a riser conversion zone to form a suspension thereof and reaction conditions maintained therein are 30 selected to particularly achieve conversion of the feed to higher yields of more desirable lower boiling products in combination with producing a product gas of relatively

high hydrogen to CO ratio. Following hydrocarbon feed traverse of the riser reaction zone in a time frame within the range of 0.5 to about 4 seconds under primarily endothermic reaction conditions to provide a riser outlet temperature within the range of 950 to 621°C 5 (1150°F), gasiform material comprising hydrocarbon vapors separated from entrained catalyst particles and separated catalyst particles are collected in the lower portion of a catalyst collecting zone 12 about the upper end of the riser discharge. A fluidizing gas not shown 10 may be charged to the bottom of a bed of collected catalyst in zone 12 to maintain it in a fluid like condition and effect some stripping of any vaporizable and strippable material from the collected catalyst. temperature high or co_2 with steam, Stripping 15 noncombustion supporting flue gas may be employed in this initial stripping of the catalyst in zone 12. Although shown, it is contemplated employing a separate stripping zone adjacent to and below vessel 12 but in for effecting initial communication therewith 20 stripping of collected catalyst particles.

It is also contemplated providing a high temperature stripping operation between the collected catalyst in zone 12 and a first stage of catalyst regeneration yet to be discussed. The high temperature stripping operation contemplated is one which mixes hot regenerated catalyst particles with spent catalyst particles in a contact zone as described in patent applications to provide a mix temperature thereof in the range of 538°C (1000°F) to about 815°C (1500°F) and contacting the catalyst so mixed with a stripping medium selected from the group consisting of high temperature

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steam, $\rm CO_2$, non combustion supporting regeneration flue gas comprising $\rm CO+\rm CO_2$, steam+ $\rm CO_2$, steam+ $\rm CO_2$, steam+ $\rm CO_2$. Generally the high temperature stripping operation will be accomplished at a temperature within the range of about 593°C (1100°F) up to about 760°C (1400°F).

The catalyst stripped by one or a combination of those herein discussed is passed by conduit means represented by conduit 14 of the drawing to a first stage of catalyst regeneration comprising bed 16 in the upper portion of vessel or regeneration zone 18 comprising fluid catalyst bed 20. The upper and lower regeneration zones are separated by pervious baffle or grid means through which regeneration gases may flow as herein after discussed.

In the first regeneration zone comprising fluid bed 16, the spent stripped catalyst particles comprising 20 deactivating deposits of hydrocarbon material hydrocarbon conversion are removed in substantial measure within the range of 40 to 80 weight percent by contact with a hot flue gas product up to about 760°C (1400°F) of the second stage of catalyst regeneration and comprising steam, CO and oxygen containing gas 25 substantially exceeding that of air. Thus the first stage of catalyst regeneration is which accomplished substantial one removal of hydrocarbonaceous deposits by the combination of endothermic and exothermic reactions in the presence of Ni, V and Fe, oxides and promoted by the reactions of 30 ${\rm CO}$ and ${\rm CO}_2$ with carbon, ${\rm CO}_2$ with hydrogen, and oxygen with carbon and hydrogen in the presence of steam (water vapor) under temperature conditions not substantially

above 760°C (1400°F) and providing a CO rich flue gas comprising steam, hydrogen, CO₂, sulfur and nitrogen oxides. Preferably CO₂ in the flue gas is kept at a low level with nitrogen oxides kept at an even lower level by employing a high purity oxygen containing gas as herein discussed. The flue gas products of the first stage of regeneration are withdrawn from zone 18 by conduit 22 for use as more fully discussed below.

Catalyst partially regenerated in bed 16 as 10 above discussed in withdrawn by conduit 24 and discharged in bed 20. In catalyst bed 20, the partially regenerated catalyst containing residual carbonaceous material or residual coke is contacted with a steam-oxygen mixture of substantially higher oxygen purity than obtainable with air and in an amount at least equal to that required to burn hydrocarbonaceous material charged to the first stage of catalyst regeneration. The regeneration gas thus described is charged by conduit 26 to a bottom portion of catalyst bed 20 to effect removal of residual 20 coke or carbonaceous material by burning in the presence of excess oxygen producing a flue gas product thereof comprising steam, oxygen and CO2 thereby heating the catalyst to an upper temperature not exceeding about A water condensation product of the 760°C (1400°F). 25 process obtained as hereafter discussed with some sulfur oxides therin is charged by conduit 28 for admixture with oxygen rich gas as herein provided. Catalyst regenerated described and comprising oxidized contaminants of Ni, V, Fe and Cu depending on feed source 30 and at a desired elevated temperature are withdrawn from bed 20 by conduit 6 for return to the lower portion of riser reactor 4 and use as above discussed.

The flue gas combustion products of the second stage of regeneration and comprising steam, oxygen and CO₂ pass from the dispersed phase of catalyst above bed 20 through a perforated grid or baffle member 30 and with the bottom portion of bed 16 for regeneration of catalyst therein as above discussed.

A CO rich flue gas comprising steam and non amounts of oxygen, combustion supporting withdrawn by conduit 22 may be passed all or in part by conduit 10 to a bottom portion of riser reactor 4 or all or a portion thereof in conduit 32 may be mixed with the gasiform product comprising hydrocarbon vapors of the separated from cracking operation riser particles and withdrawn from zone 12 by conduit 30. gasiform product in conduit 30 with or without flue gas products in conduit 32 are passed by conduit 34 to a rough separation zone 36.

In separation zone 36 a rough separation is made between C2 minus product gaseous component withdrawn by conduit 38 C₂ plus hydrocarbon components withdrawn by conduit 40, and a water phase comprising absorbed sulfur and nitrogen compounds withdrawn by conduit 42. hydrocarbon product in conduit 40 is passed fractionation equipment not shown to separate and remove gasoline boiling range material from higher and lower boiling hydrocarbon component materials. The water phase in conduit 42 is passed to a treating zone 44 wherein sulfur and nitrogen compounds are separated from the water phase and removed therefrom by conduit 46. Water thus treated is then withdrawn by conduit 48 for passage to the riser reactor 4 by conduit 8q to the regeneration

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operation by conduit 28 or to both of these operations as required and/or desired. Also a portion of this condensed and treated water may be withdrawn by conduit 49 for passage to shift reactor 56 discussed below.

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It will thus be recognized that the hydrocarbon conversion operation and the catalyst regeneration operation are in reactant balance with one another to particularly achieve upgrading of heavy hydrocarbon materials such as reduced crudes in cooperation with particularly reducing the formation of nitrogen oxides and discharge of such with sulfur oxides in the stock gas of a catalyst regeneration operation.

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The C_2 minus gas phase recovered from separator 36 by conduit 38 following separation of sulfur and nitrogen oxides therefrom in equipment not shown is passed by conduit 50 in a preferred embodiment to a methanation zone 52 wherein the C_2 minus gaseous material comprising hydrogen and CO is particularly synthesized to methane under exothermic reaction conditions. A methane rich product gas is recovered from methanation zone 52 by conduit 54 for use in providing substitute national gas.

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It is further contemplated in an alternative embodiment of passing all or a portion of the product gas in conduit 38 admixed with water in conduit 39 and/or obtained from 49 to a water gas shift operation in zone 56. A product gas rich in hydrogen and comprising CO₂ with some (CO) carbon monoxide is recovered from zone 56 by conduit 58. This product gas may also be charged directly to riser reactor 4 by conduit means not shown for use as discussed above. On the other hand, the water

gas shift product of reactor 56 is preferably passed by conduit 60 to conduit 50 and thence to the methanation zone 52 for conversion thereof to methane for utilization as suggested above.

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The alternative operating concepts of this invention are synergistically related with respect to one another to particularly suppress dehydrogenation and carbonization reactions promoted by reduced metal contaminants on the catalyst from reduced crude conversion in combinationw tih promoting hydrogenation of hydrogenation of hydrogen deficient components of the crude feedreduced This selective hydrocarbon conversion operation is implemented by the selective regeneration operation of this invention to the extent that deposited hydrocarbonaceous material are converted the regeneration operation to product particularly useful in promoting the above desired hydrocarbon conversion operation. It will be recognized by those skilled in the art that the combination of embodiments herein discussed operating contribute measurably to the economic and efficient conversion of high boiling hydrocarbons and particularly those of high Conradson carbon producing materials. The high boiling hydrocarbons particularly suitable as high boiling feeds comprise reduced crudes, vacuum tower bottoms, resids, topped crudes and synthetic oil products of coal, shale and tar sands.

In another embodiment, CO-rich flue gas from the regenerator can be recycled to the riser cracking zone so that metal-on-catalyst will act as a hydrogenation catalyst to enhance the reaction between

hydrogen produced as by-product in the riser cracking zone and CO from the flue gas. The result utilizes the CO-rich flue gas a "getter" to reduce undesirable hydrogen formation ("gassing") in the product stream exiting from the reactor.

In other modifications, the invention may be described as follows: a reduced crude of 2, or even 4, Conradson carbon number is converted into high octane gasoline and other transportation fuels with excess heat released because of the high carbon levels of the feedstock. Excess carbon is converted by the invention to CO and CO, to balance the heat released. Excess CO is removed from the regenerator, purified and combined with hydrogen produced in the riser reactor due to the hydrogenation catalyst activity of the high metals which accumulate on the cracking catalyst because of the high content of the reduced crude feedstock. resulting synthesis gas stream can be either a) converted to methane, methanol or Fischer-Tropsch products or b) using oxygen-plus-steam in place of air as the feed to the regenerator, a CO-rich product is produced which can be combined with H_2 from the cracking operation to form methane, etc. Alternatively, oxygen and steam can be fed to the regenerator to yield CO plus H_{2} so that the regenerator acts as a gasifier. As the CO/CO2-containing flue gas additionally contains SO₂ it can be recycled to the reactor as a heat source and additionally to convert SO_2 to H_2S for feed to a Claus Process sulfur removal system and the CO plus H_2 can yield CH_3 plus H_2 O (Towne gas) for fuel.

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As still an additional advantage, the ${\rm CO/CO}_2$ gas helps keep the metals on catalyst in the oxidized state to deter additional coke production.

- Lastly, hydrogen and carbon monoxide from the reactor can be combined with hydrogen and carbon monoxide from the regenerator and sent to a gas concentration unit to form methane.
- Having thus generally discussed the processing concepts of this invention and particularly described specific embodiments in support thereof, it is to be understood that no undue restrictions are to be imposed by reasons thereof except as defined by the following claims.

CLAIMS

- 1. In a reduced crude conversion process for heavy oil feeds having Conradson carbon numbers above 2, which process comprises contacting such heavy oil feeds with catalysts to form products comprising lower molecular weight hydrocarbons and coke on catalyst, said coke containing minor amounts of hydrogen and thereafter regenerating said catalyst by removing at least a portion of said coke; the improvement which comprises:
 - (a) converting said heavy oil feed in a progressive flow riser conversion zone in the presence of water condensation product separated from hydrocarbon products of said conversion operation and regenerating said catalyst in the presence of steam and CO₂ and gas containing substantial amounts of oxygen so as to produce a CO-rich flue gas,
 - (b) recovering hydrogen produced from said hydrocarbon conversion operation, separately from the hydrocarbon products of said hydrocarbon conversion operation, and
 - (c) subjecting said hydrogen to purification and removing sulfur from said CO-rich flue gas, then combining said hydrogen and said flue gas to produce a fuel gas or a synthesis gas suitable for production of methane, methanol or suitable as a feedstock for Fischer-Tropsch synthesis or high octane gasoline.

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- 2. In a catalytic cracking process for heavy oil feeds containing relatively high amounts of Conradsoncarbon-contributing materials comprising contacting such heavy oil feeds with a catalyst to form products comprising lower molecular weight hydrocarbons and coke on catalyst, said coke containing minor amounts of hydrogen and thereafter regenerating said catalyst by removing at least a portion of said coke; the improvement comprises:
 - (a) converting said heavy oil feed in a progressive flow riser conversion zone in the presence of H₂O and in the further presence of a flue gas product from said catalyst regeneration step, said flue gas product comprising primarily CO, minor amounts of CO₂ and excess steam from said regeneration operation, regenerating said catalyst in the presence of steam and CO₂ and gas containing substantial amounts of oxygen,
 - recovering a C-2 product of said hydrocarbon (b) conversion operation separate from higher boiling hydrocarbon products οf said hydrocarbon conversion operation said higher products boiling hydrocarbon comprising hydrogen and CO, and
- (c) subjecting at least a part of said C_2 product to water gas shift reaction to produce hydrogen and CO_2 .

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- In a process for riser cracking of carbo-metallic 3. oil feeds in the presence of catalyst to produce molecular lower products comprising catalyst comprising coke with and hydrocarbons subsequent regeneration of said catalyst by removing 5 said coke at high temperatures in a regeneration zone, the improvement comprising feeding H₂O and a gas comprising at least about fifty percent by volume of oxygen whereby the products of said regeneration zone consists essentially of catalysts 10 in coke content, hydrogen content, οf minor amounts monoxide with carbon dioxide, nitrogen and other gases.
- The process according to Claim 3 wherein said 4. 15 regeneration zone consists of a first regeneration stage and a second regeneration stage and wherein said coke catalyst enters said first stage and is contacted with H2O and minor amounts of oxygen and wherein said temperature is controlled to be just 20 sufficient to support the endothermic removal of coke and wherein said catalyst subsequently enters said second stage of said regeneration zone and in said second stage is contacted with substantial additional quantities of oxygen to produce catalyst 25 substantial substantially reduced coke and in quantities of CO2 and wherein the products from said first stage, consisting essentially of hydrogen, CO, minor amounts of CO2, minor amounts nitrogen and minor amounts of other gases are kept separate from 30 the gaseous products from said second regeneration stage.

- 5. In a catalytic process for cracking heavy oil feeds of relatively high Conradson carbon-contributing materials in combination with catalyst regeneration operation for removing deposited hydrocarbonaceous products of heavy oil conversion, the improvement which comprises:
- (a) converting heavy oil feed in a riser conversion zone in the presence of water condensation product separated from hydrocarbon products of said riser conversion in combination with a flue gas product of catalyst regeneration comprising primarily CO, very little CO₂ and excess steam used in the regeneration operation to achieve endothermic conversion removal of hydrocarbonaceous deposits from the catalyst in the presence of CO₂ and high purity oxygen containing gas,
- 20 (b) recovering a C₂ minus product of said hydrocarbon conversion operation separate from higher boiling hydrocarbon products comprising hydrogen and CO, subjecting said C₂ minus product of water gas shift reaction to produce hydrogen and CO₂, and
 - (c) passing the product of said water gas shift reaction to said riser conversion zone.
- 30 6. A method for upgrading high boiling oil feed materials boiling above 343°C (650°F) of high Conradson carbon level which comprises,

- catalytically converting said high boiling oil feed in the presence of a water condensation product of the process and a flue gas product of catalyst regeneration comprising primary CO and excess steam of the catalyst regeneration operation under elevated temperature conditions in a riser cracking zone and a contact time metals contaminants selected maintain to primarily in an oxidized state whereby the the operation selectivity of product improved,
- (b) -separating the products of the riser conversion operation after removal of catalyst particles comprising carbonaceous deposits into a light 15 phase comprising hydrogen, CO and gaseous methane, regenerating catalyst particles with an oxygen rich gas stream admixed with steam to produce a CO rich flue gas comprising steam, and, charging at least a portion of the CO rich 20 said riser to steam comprising gas conversion operation.
- 7. The method of Claim 6 wherein the water condensation product is separated from the product of hydrocarbon conversion comprises absorbed sulfur and nitrogen compounds, and said sulfur and nitrogen compounds are separated from said water condensation product before recycling the water to the riser conversion zone, and wherein the separated light gaseous phase comprising hydrogen and CO is subjected to catalytic methanation and the methane product thereof is used as a fuel support for the process, and wherein the

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separated light gaseous phase is subjected to water gas shift to increase the hydrogen to CO ratio of the stream before effecting methanation thereof, and wherein catalyst particles comprising carbonaceous deposits separated from hydrocarbon product vapors is subjected to a high temperature stripping operation in admixture with hot freshly regenerated catalyst before subjecting said catalyst mixture to regeneration in the presence of CO₂ rich flue gas comprising steam and oxygen.

The method of Claim 7 wherein the high temperature 8. steam operation is at least 593°C (1100°F) and stripping is accomplished with one of steam, CO_2 , or 15 mixture thereof, and wherein the catalyst recovered from said second regeneration operation is at a temperature not substantially above about 760°C (1400°F) and is stripped with an inert non-reducing gas to remove high purity absorbed oxygen therefrom before transfer to said riser conversion zone, and 20 wherein the oxygen purity of the regeneration gas stream exceeds that provided by air to reduce the nitrogen content of flue gas obtained regeneration operation and the oxygen requirements 25 combination regeneration operation initially charged to the second stage of catalyst regeneration reserved for removing residual carbon from partially regenerated catalyst, and wherein the first stage of regeneration effected with hot flue 30 gases comprising CO2, steam and high purity oxygen combination upon of endothermic exothermic reactions between CO2 and C, CO and H2O, C and H₂O and oxygen and carbon to achieve up to 80

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weight percent removal of deposited carbonaceous material and produce a CO rich flue gas free of combustion supporting amounts of oxygen.

- 5 9. A method for converting a reduced crude oil feed comprising vacuum tower bottoms which comprises, forming a suspension comprising a reduced crude oil feed, hot regenerated catalyst particles comprising:
- (a) oxidized metal contaminants, water and a CO 10 rich flue gas product of catalyst regeneration comprising water and sulfur oxides in a riser reaction zone, passing said suspension through said riser reaction zone, under conversion velocity providing conditions at а 15 hydrocarbon residence time in the range of 0.5 to 2 seconds and a suspension temperature at the riser outlet within the range of 538°C up to 621°C (1000°F up to 1150°F),
- (b) separating the suspension into a catalyst phase, a light gaseous product phase comprising hydrogen, a C₃ plus hydrocarbon phase and a water phase comprising sulfur and nitrogen compounds, and
 - (c) recycling a portion of said water phase to said riser zone following removal of sulfur and nitrogen compounds therefrom.
 - 10. The method of Claim 9 wherein the separated catalyst phase comprises hydrocarbonaceous deposits and said hydrocarbonaceous deposits are partially removed

with a regeneration gas mixture comprising CO2, steam and oxygen rich gas to produce a CO rich flue gas product comprising substantial steam in the absence of a combustion supporting amount of oxygen passed to said riser reaction zone, and wherein the separated catalyst phase is stripped temperature of at least 593°C (1100°F) in admixture with hot regenerated catalyst particles before passage to the catalyst regeneration operation and wherein the partially regenerated catalyst further regenerated to provide a high temperature regenerated catalyst of low residual coke in a separate zone with a steam-high purity oxygen containing regeneration gas steam under conditions to produce a CO2 rich flue gas comprising steam and oxygen, which flue gas is relied upon to effect said partial regeneration of said separated catalyst phase, and wherein the high temperature regenerated catalyst product of said further regeneration is recycled to each of said high temperature stripping operation and said riser reaction zone.

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